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の発明の名称

新規なペプチド及びアンジオテンシン変換酵素阻害剤

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1.発明の名称 試扱なペプチド及びアンジオデンシンの施設を指定器

2.保許益太の範囲

1) 下記の式

Lov - Lys - Pro

で思されるペプテドなよびその数。

2) 下記の式

Lau - Lys - Pro

で支されるペプテドミたはその塩を有効を分 とするアンツォテンシン要換解系退害剂。

3.美質の詳細な必須

【密葉上の利用分野】

本元朝は馬提なペプテドおよび製ペプチドまたはその覧を背頭は分とするフンジオテンシン 変技政策の監察に関する。

(延来の任事)

加圧上昇をもれらす代表的な主体内国子としてレニン・アンジオテンシン系が、また血圧原 下に困く代表的な生体内因子としてカリクレイ ントマニン平が知られているが、アンジャテンシン変後酸数(以後「ACE」という)はこのいずれの系にも大きく関与している。

一方、カリクレイン・ベニン系では四中の町 蟹体リンパク質であるのコノーデンに 四中群駅 のカリクレインが作用してチェンを速聴反びす るが、このカニンけ末梢血管を拡張させるとと

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もにホス本リパーゼムを新性化してプロステア ラッンの合成を関盤して血圧を降下させる。と ころがこのカリケレイン・キニン系にACSが 個くと、ACSは東側血管の感染作用およびポ ユテリパーゼムの唇性化作用を含する上記をニ ンを分解・水低性化してしまうために、血圧の 降下が坐じなくなる。

したがって、ACEの上記のような個をも個書する物質(ACE観告組)が存在すると、血匠上外的質であるアンジャインンとの生成が抑制され、及つ原圧降下物質として降くキュンの分離が防止されて、血症の上外内割割よび血圧性でが可感になる。

かから点がり世年人でも担告別の研究問題が 色っ行われており、犬はダンパク質由来の、ま たは合成による特定のペプチド類が人でを阻害 作用を育することが経色されている。これまで に特容された天然タンパグ質由来の人でを観言 ペプチドとしては、マムシ白米のブラジをニ ン、ボテンシューターを(Pyr-Ciy-Las-Fra-Pro

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その必要、上記既知のACを観察ペプチドとは又なったアミノ既配列を有する、ロイシンーリジンープロリンが配列した初まなトリペプチド lea-lya-Pro を気波ランパク質の加水分解物から年載することができ、そしてこのトリペプチドがACを観音作用を有することを具置した。

したがって、本発別は、下型の丈 Leu-Lys-Pro

で表されるペプチドおよびそのはである。 更に、本発明は上記式で表されるペプテドも たはその生を自然成分とするACE組名割を包 きする。

本生初の上記ららなほお商品を育するベブテドは、最初は乳器タンパリ質のプロテアーゼによる加水分別処理生産物をして発見されたものであり、その場合には上記3番のアミノ酸である。 LysなよびProはいずれもL=アミノ酸である。 しかしながら、それに設定されず上記のアミノ 配配列を有するトリベブをどであればいずれの -Brg-Pra-Lye-He-Pra-Pra)およびプラツギニ ン・ポテンシェーダーで (Pyr・Cly=Leu-Pro-Pro -Gly-Pro-Pro-11e-Pro-Pro) (いずれる H. Saco and T. Suzukl. Brochemistry. 10. p.972 (1971) 瓜記載されている)、 年見カゼイン出来 のペプナドである Pha-Pac-Val-Sis-Pro-Pho-Pro-Glu-Fal-Phe-Gly-Lya (森 公 68 60 - 23085年 全型)、Pho-Phe-Yal·Ala·Pro (好明 昭 59·44323 今公禄〉、Thy-Thr-Mec-Pro-Leu-Tro(神四平3-20283年公司)、Ala-Yal-Pro-Tsr-Pro-Gin-Ara. 魚菜タンパク質由素のペプチドであるTyr・Lrs− Ser-Pac-lie-Lyg-Gly-Tyr-Pro-Val-Wet, Pro-Clu-Glu-Glu-Pro-His-Val·Leu、トクモロコシ ァーゼイン由来のペプラドである Leu-Pro-Pro. Val-His-Len-Pro-Pro. Val-His-Leu-Pro-Pro-Pro 中を思げることができるが、その大手は? ミノロが5便以上取合したペプナドである。 (長明の内容)

上記のような状況下にお妻褒者らもACE鼠 雲な月を有する動質について同名を進めてきた。

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犬字器性なであってもよく、なう思クアミノなの全部がローアミノ版からなるトリペプチドなよびう音のブミノ版のうちのいずれかしつまたは2つがレーアミノ酸であって無ながローアミノ種からなるトリペプチドも包含され、それらは化学会成によう製造することができる。

本発明のトリベブナドの顧其法の例を挙げる と以下のとおりである。

乳オクンパク質の四水分解による方法

利請タンパク質をプロチアーでを使用して加水分解して水影性の乳情タンパク質由来ペプチド配合物を調整する。その際に、乳管タンパク質を水管の数体中に分離をたは経路をせた状態で加水分解を行うのが、治疗のし品さ、目的物の収益の範囲の床から分ましい。

プロテアーゼとしては、監性で作用するプロ ナアーゼ、特に終業の監性中心にアスパテモン 型達玉とアスパラギン細のカルボン酸イポンが 関手するアスパルティックプロテイナーゼを運 用するのがよい。そのようなプロテアーゼの例

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としては、ペプシン、ヒイログゲ母素のアスパ ルティックプロティナーゼ、アスペルダルス型 森のアスパルティックプロティナーゼ。 ペニシ リウン武型のアルバルティックプロティナーゼ を平けることがです。背にペプシン。アスペル チルス返尿のアスパルティッシプロティナーゼ しい。プロテアーゼは1銭短のみを使用しても、 またはプロテアーゼ日士がお互いた悪影響を及 ほさないかざりは装数値を併用してもよい。在 数のプロチナーゼを使用する場合は、超延数の プロチアーゼも国際に非正させて即水分裂を 打っても、またはく着菜でつ選びに用いて加木 分界を行ってもよい。また、プロテアーゼはフ リーの状態で使用しても、固定让レイ控用して もよい。プロテアーゼの奴別並はいずれの語合 も更盛したダルテン|00ま当たりプロテアーゼ 約5.000~100.000 malcat用いるのがよい。

ここで本明元を中のブロテアービ新姓(eAit) はすべて下記の方式により測器したものであ

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タ質等の不溶性変形的を減心分配のの過当な手 色で分解数差し、強密性中に全まれているペプ チド混合物を乾燥等により固数する。

次いて、このペプテドを含物を水等に根壁を せた状態で配分面、イボン交換分面、ゲル等過 分置等により分解符製し、それを更に高速度体 クロマトグラフィー(例えば運用カラムを用い た高速度体クロマトグラフィー等)等により処 をして上記とリペプチドを純粋な形容で無風する。

上売したペプチドス合物を含育する水の取の 分配額整およびトリペアテドの単題は、作えば 次の(s)ー(1)の工器かりなる方法で行うことが てまる。

(a) ペプチド語の物を含ぎする本容度のpile 的 3.0~5.0に関数し、これをイオン交換クロヤトダファィーにかけ(存えば取り一株式会社 なの SP-Tayopears 550 C を見扱したカラムに 高速させる)、このクロマトダラフィーに要 含しに収分を O M から0.5 M までの道線速度

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プロマアーゼ反性の無国法

監督として末旬ノルク社型のハマーステインカゼイン1%直接を用い、アンソン一級原理性 (連絡の感染 "野裏研究法" 中2 年,第237页(略和30年1月10日,終身委囚犯者)] により加定した。反応は30でで30分配分い、1分間に148のテロシン組出量を避験するのに減ずる解釈無を1unitとした。

プロテアーゼン図は、ちゃの状況(例人は プロケアーゼの温度、プロテアーゼの使用を感 書)に応じて最適のpH、温度、プロテアーセ金、 出間温度、処理時間手の気体を温度して行うの がよく、例えば上で挙げたプロテアーゼを放斥 する場合にはpH的1.5~5.0、四度四30~50でで、 0.75以トリクマロ函数への利用率が約40~70% になるまで知水分類を行うとよい。

自的とするMIX分離共感が選成された時点で 加熱および/またはoH 温度してプロチェーゼ を実施させ、免済した野豚、未分解乳湯タンパ

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为配を含するNaCi水町級でお除し、得られる名部分の中から高い服容は性を育する部分(Maci水町鉄路景が約0.4~0.5 Mの箱頭で浴板してくる部分) を回収する。

- (b) 上記高い曜舎を使き合する百分と分子あるい 処理して (例えばパイナタッド社業のパイオゲルアー2 を充填したカラムを過過させる)、更にいくつかの西方に数の水でぬかかせしてその少から更に高い温名を使そ者する 異分も回収する、
- (c) 上記(b) で国民した日介を高温版学リロマトグラフィー (例えば近ソー株大会装品の ODS ー 120T) に通過させ、受参成分を0.1%トリフルまの時度水母級 (A 版) とアセトニトリルを50%会有する0.1%トリフルギロ部飲水母級 (B 版) との資金費であって配合性のの各種の監証が0%から100%をで国際的に増加する配金費を到配用販売を用いて搭載すると、アセトニトリルの書座が約20-22%の範囲の母組版区分に英級など-2が現れ、この

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母会のACE母官 性に創生事題して四及する。

- (d) 必要に応じて上名(c)の工程を線返す。(a) (d)工程で得られた配分から必需を電道等に より映金して自色の関係を類なし、そして
- (1) 上記白色圏体として移られた生成物のアミン酸配列を例えば島障視が原製の気根式プロティンショブンサー(PSQ= l ソステム) 等を を風して遊べ、Leu-Lys=Proからなるより パプナドであることを確認する。

また、本発酵のトリペプチドを化学会配により製造する場合は、例えば次の方法を経用することができる。

水苑明の)リペプチドの化学会系基

ペプチド台成英配(ファルマンア社(スエーデン)製のBiolonx 4176)を使用して合成する。 具体的には、ポリアミド制度にFees・ブロリン を招合をはた後もの Face 至を放棄して必然ア ミノ蓋を透離させ、この運転アミノ蓋にFees・ リクンを組合させてから Face 苦を飲棄し、更

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と一括に発与してもよく、 図は他の袋割と適合または組合わせて使用することができる。 またほう影似は、 設剤、 九利、 毎粒剤、カブセル、 込刻、水波液、 任針所等の任金の母様が可能で ある。

夏に、本先男のACを贈答別は、会品や母呼中に恐かして、またほぞれらと一葉に甘みすることもでき、その場合には天教リンパク質に自来する C-lau-L-Lyu-l-Pro が登ました。

以下に、本第明をあき申げて具体的に契明するが本発明はそれのによって限定されない。 史 編 第

私法のンパグ長(日本プロタインを完全社 ALACEN 132)5gを0.03N塩酸100mgに分散回解させた後、匹留水を耐えて全量200mgにした。 1 円塩酸を加えてpHを2.0に調整した後、ペプシン(本国ングマを取)5000mitsを加え、37つで15時間反応させた。次に、5 N水酸化ナトリッム水の政でpBを4.4に調査した後、アスペルギルス起変のアルバルチェックプロティナー

C fmac・のインンを組合してからfmac易を飲金してお望い路で保護されたペプテドリ形成する。これを95%トリフルキの計画水構成と生意で60分類反応させてお聞き分成させた後、 明期を調かする。トリフルキの配像水格減を設置を示した後、 受害を20.1 N配給に合同し、その合成を高速値体グロマトグラフィー(008-180T)に対して不同的を収益することによって関係の高い igu-154-Pro を単純する。

本差別のACE監察剤は人間および包ェの動物に数多することができ、少量の数をによって 類音な魚圧降でおよび上昇和似を造成すること ができる。

本見明のACE関告制の計画なな子会は、最 子をれる人間や動物の多名、作品、性別、流状、 動物の相差年の有々の美井によって異なる。

そして、本見明のACEMを利は品口ならおよび非年の長らのいずれによっても数手可能であり、変に単独でならしても、また製造工業にないて過剰使用されている個の資源や遊び出来

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で(大野和田社製のプロテアーゼM)]000mnitsを加えて45でで5時間反応させた。次いで、5N水銀にアトリウム水気機でpHを6.0に減空した後90でで20分間加熱して酵素を大利させるとよらに未給物を記録させた。田田にお知した後、10000cで20分間減心分離して固か物を分配除去した。上陸液を関取して環府必然してペプクである毎4.0sを得た。

上記で終たペプチド記名数500mgを5mm的政 最低級50mgに対象した後、 | N塩酸でpH3.5に 減型した。

これを選逐15mm、長さ200mmのカラムに置い 一体式会社器のSP-Toyopearl 550Cを40m2光 ほしたイナン交換タロマトカラムに1.0m2/分 の建建で冷温させた後、このクロマトカラムに 気器した成分もりMから0.5Mまでの医総合器 お配を有するPaci本間級からのも密節器 120ms を1m4/分の設定で返してカラムかの研覧した ところ、Naci水形蔵を繋が0.4~0.5Mの部分に 高い相等を性を買する置分を終たのでこれを図

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女した。

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表かて、上記画分をパイナテッド社製のバイオゲルP = でも200m8充切したカラム(カラム既通16mm、長さ1000mm) にも.33mm/分の改変で過して分子からい処理し、まに重要水で搭降してその中から高い理管居住を有する国分を図収した。

得られた西分から本質を発換施設して自己の

-- IS -

をせる。次に3 N 総数2000c M 元、 数四水で50 恰に出まする。約30分数に分元数元元を元式で砂 起表兵300pm、 気光波及490pmにおける生光強度 (A)を固定する。 解析状の代わりにな留を50mc を同様に外国して出光処皮(B)を制定する。

過者思想はB-A/Bにより果わられる。

飲料底の機変を放えて、経営過程を上型と回答に排足し、活性を50%結合する会交を求めて これを10、として長した。

(ペプテドのACB組合连佐(ICLa))

<u> </u>	10: (4H)
E - L-Len-L-Lys = L-Pro·CH(本発明)	2.3
ブラジキニン・ポテンツエーサーB	ō-4
ブラグギニン・ポテンシエーターC	29.0

上記器の応見から、本発明のACE組合製は 既知のACE組合物ブラジをニン・ポテンシメ ーチーリおよびCに比べて極めて低級皮質で、 マなわちごく少量の使用で「Cuvを選択するこ とができ、ACE組合版性が実際に高いことが 部は i200maを回収した。この四色図はを単常社 作所製の気料式プロティンシーケンサー (PSO-| システム) を使用してそのアミノ酸配列を細 べたとこう、ド末編から順次し-Leu、i.-l.yaがよ びi-Proが返路してきた。このことから六 H・l -leu-l-lys-l-Pro・OB であられることが在記された。

上記で開製したトリベブテドおよび収録の A C E M 空ベブチドのA C E M 要成性を下述の 方法で制定したところ、下記の正に示すは果る 併た。

ベプチドのACE別事后性の制定法

状は屋SOutを放験者にとり、これにACE板(単位シリマは取のうさぎ間由来のACEの!
unitを来ちndに合解させたもの)20Aをかえる。
37でに1分間除った後、亜質(Suk Bip-His-(Go : p88.3)を加えて37でで30分屋ださせ、犬いて0、3 M水酸化ナトリッム水溶液1 ngを加えて22でで20分屋ださせ、アンビを存止させる。外土就要オルトーフタルフルデモド約100mgで加えて22環で10分階反応

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n # 8.

【発明の効果】

本発明のACE組制的は、窓のて少量の食を でACEの活性で思想して四肥能でおよび血圧 上昇物製を選択することができる。

1た、水契明のACE四宮副は、 口色の水部 性物取であるために、 そのままでまたは水等に 砂度をマモ毎の食みおよび赤品口を手のいずれ の方法によっても極めて原当に、皮与することが できる。

そのよ、本発酵の新設なトリペプチド Lead-Lys-Pro は、3個のアミノ酸が配列しただけの変むで簡単な関連を有する能分子変化の物であるため、化学育成によっても固単に製造することができ、しかも交易した場合に体内での気収性がよく高い血圧発汗作用を示す。

特許山田人 日 神 野 郡 井 史 今 廷

化引入 非位世 四 不 于



平成

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訳

表

(平成4年6月2日発行)

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施州紀号

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出願人名称 日精製粉株式会社 (日次とも)

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NEW PEPTIDE AND ANGIOTENSIN-CONVERTING ENZYME INHIBITOR

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UNITED STATES PATENT AND TRADEMARK OFFICE
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NEW PEPTIDE AND ANGIOTENSIN-CONVERTING ENZYME INHIBITOR

[Shinkina peptide oyobi angiotensin henkan koso sogaizai]

Inventor:

Hirofumi Motoi

Applicant:

Nissei Seifun K.K.

[There is no amendment in this translation]

Claim

1. Peptide and its salt expressed by the following equation.

Leu-Lys-Pro

2. Angiotensin-converting enzyme inhibitor having the peptide or its salt expressed by the following equation as the effective component.

Leu-Lys-Pro

Detailed explanation of the invention

Industrial application field

This invention concerns a new peptide and angiotensin-converting enzyme inhibitor having said peptide or its salt as the effective component.

Prior art

The renin-angiotensin system as a representative in vivo factor which results in raising blood pressure and the kallikrein-kinin system as a representative in vivo factor which functions to lower blood pressure are known, and the angiotensin-converting enzyme (will be referred to as "ACE" below) contributes greatly to both of these systems.

The mechanism will be briefly explained. First, in the renin-angiotensin system, the enzyme renin from the kidney which is secreted into the blood interacts with angiotensinogen in the blood, and forms angiotensin I, which is a decapeptide. This angiotensin I does not display the blood pressure raising action; however, when ACE interacts with this, it forms angiotensin II, which is an octapeptide. This angiotensin II contracts the peripheral blood vessels and also interacts with the adrenal cortex and promotes the production of aldosterone. Aldosterone interacts with the kidney and results in an increase in the heartbeat output rate through the invitation of the readsorption of sodium and an increase in the amount of body fluid. Both of them significantly increase the blood pressure.

On the other hand, in the kallikrein kinin group, the kallikrein as the enzyme in the blood interacts with kininogen, which is a precursor protein in the blood, and frees and produces kinin. However, this kinin dilates the peripheral blood vessels and also activates phospholipase A₂, promotes the synthesis of prostaglandins, and lowers blood pressure. However when ACE interacts with this kallikrein-kinin group, ACE breaks down and inactivates the aforementioned kinin, which has the dilating action of the peripheral blood vessels and the activation action of phospholipase A₂, and lowering of blood pressure does not occur.

Accordingly, when a substance which inhibits the aforementioned action by ACE (ACE inhibitor) is present, the formation of angiotensin II, which is a blood pressure raising substance, is inhibited, and the breakdown of kinin, which functions as blood pressure lowering substance, is prevented as well, and the control over raising the blood pressure and lowering the blood pressure becomes possible.

From such a viewpoint, various types of research and development of ACE inhibitors have been implemented in recent years, and it has been reported that specific peptides that originate from natural proteins or through synthesis have the ACE inhibiting action. The ACE inhibitors originating from natural proteins that have been reported so far include bradykinin potentiator B (Pyr-Gly-Leu-Pro-Pro-Arg-Pro-Lys-Ile-Pro-Pro) and bradykinin potentiator C

(Pyr-Gly-Leu-Pro-Pro-Gly-Pro-Pro-Ile-Pro-Pro) that originate from the mamushi (both are described in H. Kato and T. Suzuki, Biochemistry, 10, p. 972 (1971)], milk casein originating peptides Phe-Phe-Val-Ala-Pro-Phe-Pro-Glu-Val-Phe-Gly-Lys (Japanese Kokoku Patent No. Sho 60[1985]-23085), Phe-Phe-Val-Ala-Pro (Japanese Kokai Patent No. Sho 59[1984]-44323), Thr-Thr-Met-Pro-Leu-Trp (Japanese Kokai Patent No. Hei 2[1990]-20263), Ala-Val-Pro-Tyr-Pro-Gln-Arg, fish protein originating peptides Tyr-Lys-Ser-Phe-Ile-Lys-Gly-Tyr-Pro-Val-Met and Pro-Glu-Glu-Glu-Pro-His-Val-Leu, corn

Tyr-Lys-Ser-Phe-Ile-Lys-Gly-Tyr-Pro-Val-Met and Pro-Glu-Glu-Glu-Pro-His-Val-Leu, corn γ-zein originating peptides Leu-Pro-Pro, Val-His-Leu-Pro-Pro, and Val-His-Leu-Pro-Pro-Pro, for example. The majority of these are peptides in which 5 or more amino acids are linked together.

Content of the invention

Under the aforementioned circumstances, the inventors of this invention also have advanced the research on the substances having the ACE inhibiting action.

As a result, it has been discovered that a new tripeptide Leu-Lys-Pro, in which leucine-lysine-proline are arranged with an amino acid sequence different from the aforementioned existing ACE inhibiting peptide, can be isolated from a hydrolyzed milk serum protein, and this tripeptide has the ACE inhibiting action.

Accordingly, this invention is a peptide and its salts expressed by the following equation. Leu-Lys-Pro

This invention also includes the ACE inhibitor with the peptide or its salt expressed by the aforementioned equation as the effective component.

The peptide having the aforementioned ACE inhibiting action in this invention was first discovered as a product through a hydrolysis treatment by the protease of a milk serum protein, in which all of the aforementioned 3 kinds of amino acids Leu, Lys, and Pro are L-amino acids in that case. However, without being limited only to these, any optical isomer can be used if it is a tripeptide having the aforementioned amino acid sequence, including tripeptides in which all of said 3 types of amino acids are made of D-amino acids, and tripeptides, in which any one or 2 of the 3 types of amino acids are L-amino acid and the rest are D-amino acid, and they can be manufactured by chemical synthesis.

An example of the preparation method of the tripeptide in this invention is listed below.

Method by the hydrolysis of a milk serum protein

A milk serum protein is hydrolyzed by using a protease, and a water-soluble milk serum protein originating peptide mixture was prepared. During this, a hydrolysis of the milk serum protein in a condition in which it was dispersed or dissolved in a liquid, such as water, for

example, is desirable from the viewpoints of easy operation, yield of the target product, and purity.

As the protease, a protease that reacts in acid is desirable, and, the use of an aspartic protease having the aspartic acid residue and the carboxylic acid ion of the aspartic acid related to the activity center of the enzyme is particularly desirable. Examples of such a protease include pepsin and orange polyporus-originating aspartic protease, Aspergillus-originating aspartic protease, and Penicillium originating aspartic protease. Pepsin and the Aspergillus-originating aspartic protease in particular are desirable at the point of obtaining the target product at a high yield. Only 1 kind of protease can be used, or several types can be used in combination as long as there is no negative effect extended among the proteases. When using several types of proteases, said several proteases can be simultaneously present and hydrolyzed, or 1 kind may be successively hydrolyzed at a time. The protease may be used in a free or stabilized state. The ideal amount of the protease that is used in all cases is about 5000-100,000 units per 100 g of dry glutene.

The protease activity (unit) in the specifications here is entirely measured by the method below.

Measurement method of the protease activity

Using a 1% Hammerstein casein solution manufactured by US Melk Co. as the matrix, it was measured by the Anthon-Hagiwara [transliteration] Modified Method (Edited by Shiro Akabori, "Enzymatic Research Method," Vol. 2, Page. 237 (published on January 10, 1961 from Asakura Books)). The reaction was held at 30°C for 30 min, and the amount of enzyme required for isolating tyrosine in an amount equivalent to 1 µg in 1 min was established as 1 unit.

It is desirable to select the conditions for the optimal pH, temperature, protease amount, processing speed, and the processing time, etc., of the protease process according to various circumstances (for example, the type of protease, the form of protease used, etc.). When using the protease listed above, for example, a hydrolysis may be performed at a pH of about 1.5-5.0 and a temperature of about 30-50°C until the dissolution ratio into a 0.75M trichloroacetic acid reaches about 40-70%.

At the point of reaching the target state of hydrolysis, the protease was inactivated by heating and/or adjusting the pH, the inactivated enzyme and insoluble solids, such as the milk serum protein that had not decomposed, for example, were separated and eliminated by a proper measure, such as centrifugal separation, for example, and a peptide mixture contained in the residue solution was collected through drying, for example.

Successively, this peptide mixture in a condition in which it is dissolved in water, for example, was separated and purified through a membrane fraction, ion-exchange fraction, and a

gel filtration fraction, for example. This was furthermore processed through high-speed liquid chromatography (high-speed liquid chromatography using a reverse-phase column, for example), for example, and the aforementioned tripeptide was isolated in a pure form.

The separation and purification of an aqueous solution containing the aforementioned peptide mixture and the isolation of a tripeptide can be obtained by a method consisting of (a)-(f) processes below.

- (a) The pH of the water solution containing the peptide mixture was adjusted to about 3.0-5.0; this was run through ion-exchange chromatography (passing through a column packed with SP-Toyoperal 550 C manufactured by Toso K.K., for example), the component which had adsorbed onto this chromatography was eluted by a aqueous NaCl solution having a linear concentration gradient from 0M to 0.5M, and fractions having a high inhibition activity (fractions in which the concentration of the NaCl water solution was eluted within a range of about 0.4-0.5M) were collected from the fractions that were obtained.
- (b) The aforementioned fractions having a high inhibition activity were processed by a molecular filtration (passing through a column packed with Biogal P-2 manufactured by BioLad Co., for example), and were furthermore eluted and separated into several fractions by distilled water, and fractions having a higher inhibition activity were collected from them.
- (c) The fractions collected by the aforementioned (b) were passed through a high-speed liquid chromatography (ODS-120T manufactured by Toso K.K., for example), the components which were adsorbed were eluted by a linear concentration gradient eluent, which was a mixed solution of a 0.1% aqueous trifluoroacetic acid solution (solution A) and a 0.1% aqueous trifluoroacetic acid solution containing 50% acetonitrile (solution B), in which the concentration of solution B in the mixed solution increases linearly from 0% to 100%, a high adsorption peak appeared in the section of the eluent within a range of about 20~22% of the concentration of acetonitrile, and the ACE inhibition activity of this fraction was measured and confirmed, and collected.
 - (d) The aforementioned process (c) was repeated if necessary.
- (e) The solvent was eliminated through drying, for example, from the fraction obtained through process (d), a white solid content was collected, and
- (f) The amino acid sequence of the product obtained as the aforementioned white solid was checked by using a gas-phase protein sequencer (PSQ-I system) manufactured by Shimazu Seisakusho, for example, and the tripeptide consisting of Leu-Lys-Pro was confirmed.

Also, the following method can also be adopted, for example, when manufacturing the tripeptide in this invention through a chemical synthesis.

Chemical synthesis method of the tripeptide in this invention

It was synthesized using a peptide synthesis device (Biolynx 4170 manufactured by Pharmacia Co (Sweden)). Concretely, after condensing the Fmoc·proline with a polyamide resin, that Fmoc radical was removed, the terminal amino acid was isolated, the Fmoc·lysine was condensed with this free amino acid; then, the Fmoc radical was eliminated, and Fmoc·leucine was also condensed, the Fmoc radical was eliminated, and a peptide protected by the aforementioned resin was formed. This was reacted with a 95% aqueous trifluoroacetic acid solution at room temperature for 60 minutes and the resin was separated, and the resin was eliminated through filtering. After eliminating the trifluoroacetic acid water solution under reduced pressure, the residue was dissolved in 0.1N acetic acid, that solution was passed through a high-speed liquid chromatography (ODS-120T), impurities were eliminated, and Leu-Lys-Pro at a high purity was isolated.

The ACE inhibitor in this invention can be administered to humans and various types of animals, and a significant lowering of blood pressure and a control of its raising can be attained through a dosage in a small amount.

The satisfactory dosage of the ACE inhibitor in this invention is different according to various types of conditions, such as the age, weight, sex, and conditions of the humans and animals and the type of animal, etc., for the administration.

Then, the ACE inhibitor in this invention can be administered through both oral and nonoral administrations. Furthermore, it can be administered independently, or may be administered together with a solid support and a liquid support which are generally used in the pharmaceutical industry. Or, it can be used mixed together or combined together with other substances. Also, possible administration forms include any optional forms including tablets, round tablets, granulars, capsules, dispersing agents, aqueous solutions, and injection agents, etc.

Furthermore, the ACE inhibitor in this invention may be added into foods and feeds or can be administered together with them, and L-Leu-L-Lys-L-Pro which originates from a natural protein is ideal in that case.

This invention will be concretely explained in an example below; however, this invention should not be limited by it.

Application example

After dispersing and dissolving 5 g of milk serum protein (ALACEN 132 by Nippon Protein K.K.) in 100 mL of a 0.03N hydrochloric acid, distilled water was added to a total volume of 200 mL. After adjusting the pH to 2.0 by adding 1N hydrochloric acid, 5000 units of pepsin (manufactured by US Sigma Co.) were added, and reacted at 37°C for 15 h. Next, after adjusting the pH to 4.4 by the 5N aqueous sodium hydroxide solution, 1000 units of alpaltic

[transliteration] protease originating from Aspergillus (Protease M manufactured by Amano Pharmaceutical Co.) were added, then reacted at 45°C for 5 h. Successively, after adjusting the pH to 6.0 with the 5N aqueous sodium hydroxide solution, [the solution was] heated to 90°C for 20 min and the enzyme was inactivated, and the undissolved substances were precipitated. After cooling to room temperature, solid contents were separated and removed through a centrifugal separation of 10,000 G for 20 min. The supernatant was collected, freeze-dried, and 4.0 g of a peptide mixture were obtained.

500 mg of the aforementioned peptide mixture obtained were dissolved into 50 mL of a 5mM acetic acid buffer solution; then, the pH was adjusted to 3.5 by the 1N hydrochloric acid.

This was passed through an ion-exchange chromato-column packed with 40 mL SP-Toyopearl 550C manufactured by Toso K.K. in a column with a diameter of 16 mm and a length of 200 mm at a flow rate of 1.0 mL/min, the components that were adsorbed onto this chromato-column were eluted from the column by running 120 mL of an eluate consisting of the aqueous NaCl solution having a linear concentration gradient of 0M to 0.5M at a flow rate of 1 mL/min, fractions having a high inhibition activity in the area of 0.4-0.5M in the concentration of the aqueous NaCl solution were obtained, and they were collected.

Successively, the aforementioned fractions were passed through a column (diameter of the column is 16 mm, and the length is 1000 mm) packed with 200 mL of Biogal P-2 manufactured by BioLad Co. at a flow rate of 0.33 mL/min, processed by a molecular filtration, eluted with distilled water next, and fractions having a high inhibition activity were collected from them.

After passing the aforementioned fractions through a high-speed liquid chromatograph ODS-120T manufactured by Toso K.K. at a flow rate of 1 mL/min, the components that were adsorbed were eluted by running an eluate, which was a mixed solution of a 0.1% aqueous trifluoroacetic acid solution (solution A) and a 0.1% aqueous trifluoroacetic solution containing 50% acetonitrile (solution B), and has a linear concentration gradient in which the concentration of solution B in the mixed solution increased linearly from 0% to 100%, at a flow rate of 1 mL/min, the concentration of the acetonitrile had a high inhibition activity in the section of the eluate of 20-22%, these fractions were collected, and this high-speed liquid chromatography processing was repeated again.

The solvent was dried and eliminated from the fractions that were obtained, and 1200 µg of a white solid content were obtained. The amino acid sequence of this white solid content was checked by using a vapor phase type protein sequencer manufactured by Shimazu Seisakusho (PSQ-1 system), L-Leu, L-Lys, and L-Pro successively isolated from the N terminal. Through this, the tripeptide expressed by equation H·L-Leu-L-Lys-L-Pro·OH was confirmed.

The ACE inhibition activity of the tripeptide prepared above and the existing ACE inhibiting peptides were measured by the method below, and the results indicated in the table below were obtained.

Measurement method of the ACE inhibiting activity of peptides

 $50~\mu L$ of the sample solution were put in a test tube, $20~\mu L$ of the ACE solution (1 unit of ACE originating from a rabbit lung manufactured by US Sigma Co. was dissolved in 5 mL water) were added to this. After maintaining this at 37°C for 5 min, the substrate (5 mM Hip-His-Leu: pH 8.3) was added, and reacted at 37°C for 30 min. Successively, 1 mL of a 0.3M aqueous sodium hydroxide solution was added, and the reaction was stopped. $100~\mu L$ of a fluorescent test drug orthophthalaldehyde solution was added, and reacted at room temperature for 10 min. Next, 200~m L of 3N hydrochloric acid were added, and diluted 50 times with distilled water. The fluorescent intensity (A) at the excitation wavelength of 300 nm and the fluorescent wavelength of 490 nm were measured by a spectrofluorophotometer after about 30 min. $50~\mu L$ of distilled water were similarly processed instead of the sample solution, and the fluorescent intensity (B) was measured.

The inhibition activity can be obtained by B-A/B.

Changing the concentration of the sample solution, the inhibition activity was measured in the same aforementioned manner, the concentration for inhibiting 50% of the activity was obtained and this was expressed as $1C_{50}$.

①[ペプチドのACE阻害活性(IC)]
2 × 7 + 5	1C. (μΝ)
H·L-Leu-L-Lys-L-Pro·OH (本発明) 3	2.2 -
ブラジキニン・ポテンシエーターB4	6-4
ブラジキニン・ポテンシエーターC 5	29.0

- Key: 1 (The ACE inhibition activity $(1C_{50})$ of peptide)
 - 2 Peptide
 - 3 (This invention)
 - 4 Bradykinin potentiator B
 - 5 Bradykinin potentiator C

From the results in the aforementioned table, it can be understood that the ACE inhibitor in this invention is a solution with a very low concentration when compared to the existing ACE inhibitors bradykinin potentiators B and C, in other words, 1C₅₀ can be attained in a very small amount of use, and the ACE inhibition activity is very high.

Effect of the invention

The ACE inhibitor in this invention inhibits the ACE activity and attains a lowering of the blood pressure and a control of the raising of the blood pressure through its administration in a very small amount.

The ACE inhibitor in this invention is also in the form of a white water-soluble powder; therefore, it can be very easily administered either through oral administration or nonoral administration directly or while being dissolved in water, for example.

Moreover, the new tripeptide Leu-Lys-Pro in this invention is a low-molecular-weight compound having a very simple structure where only 3 amino acids are arranged, which can be easily manufactured through a chemical synthesis as well. Moreover, it displays an excellent adsorptivity in the body and a high-blood-pressure lowering action when administered.

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- 1. Title: New Peptide and Angiotensin converting enzyme-inhibitor
- 2. Range of Patent petition
 - 1) Peptide Leu-Lys-Pro and its salt
 - 2) Angiotensin converting enzyme-inhibitor from peptide Leu-Lys-Pro and its salt.
- 3. Description of invention

(Extent of commercial use)

This invention is about angiotensin converting enzyme-inhibitor from new peptide Leu-Lyn-Pro, existing one, and their salt.

(Former technique)

Renin-angiotensin system (r-a) is one of the factors of hypertension. Kallikrein-kinin system (k-k) is one of the factors of antihypertension. Angiotensin converting enzyme (ACE) takes part in both of them. In r-r in blood, renin from kidney produces angiotensinI, decapeptide, with angiotensinagen. When angiotensinI reacts with ACE, it produces angiotensinII that contracts peripheral blood vessel and promotes aldosterone from adrenal cortex. Aldosterone makes kidney reabsorb sodium and increase body fluid to increase blood pressure.

On the other hand, in k-k, kininogen reacts kallikrein to release kinin that expands peripheral blood vessel, activates phospholipaseA2, and accelerates synthesis of prostaglandin to reduce blood pressure.

However, when ACE reacts to r-a, ACE breaks and deactivates kinin. Therefore, if ACE inhibitor exists, it is possible to suppress producing angiotensinII and breaking kinin.

Recently some researches have indicated special peptides made from nature and synthesis inhibit ACE. Natural ACE inhibiting peptides:

-From pit viper: bradykinin potentiater B (Pyr-Gly-Leu-Pro-Pro-Arg-Pro-Lys-Ile-Pro-Pro) bradykinin potentiater C (Pyr-Gly-Leu-Pro-Pro-Gly-Pro-Pro-Ile-Pro-Pro)

(H.Kato and T.suzuki, Biochemistry, 10, P.972, 1971)

-From milk casein: Phe-Phe-Val-Ala-Pro-Phe-Pro-Glu-Val-Phe-Gly-Lys(patent pub.#6023085)

Phe-Phe-Val-Ala-Pro(patent pub.#5944323)

Thr-Thr-Met-Pro-Leu-Trp(patent pub.#0220263)

Ala-Val-Pro-Tyr-Pro-Gln-Arg

-From fish protein: Tyr-Lys-Ser-Phe-Ile-Lys-Gly-Tyr-Pro-Val-Met

Pro-Glu-Glu-Pro-His-Val-Leu

-From corn γ-zein: Leu-Pro-Pro

Val-His-Leu-Pro-Pro Val-His-Leu-Pro-Pro

Most of the peptides are more than 5 chains of amino acid.

(Content of the invention)

As the result of the inventor's research, he has invented that he can extract new peptide from hydrolyzed whey protein that inhibits ACE.

This invention is the amino acid sequence, Leu-Lys-Pro, and its salt.

This invention includes ACE inhibitor.

The ACE inhibiting peptide is found after hydrolysis of whey protein with proteinase. These 3 amino acids are all L-Amino acids, but optical isomer is not matter if tripepetide that has all 3 amino acid is used.

These amino acids can be even produced by synthesis from tripeptide of D-amino acid that has one or two L-amino acid.

The materials for hydrolyzing whey protein

- -Whey protein solution w/distilled water
- -Proteinase for acid condition

Aspartic proteinase: from pepsin; Aspergillus; Penicillium.

Best if use one from pepsin or Aspergillus.

The proteinase can be mixed with others before and after adding to the whey solution.

The condition of the proteinase is ether free or fixed.

The concentration of the proteinase5,000~100,000 units per 100g gluten powder.

The method of measuring proteinase activity

- -Use 1% solution of hammer stain casein (Melk, U.S.)
- -Use Anson-Hagiwara Method (S. Akahori, "Enzyme Research Method", No.2, p. 237, 1/10/1961, Asakura Book)
- -1 unit=the amount of enzyme to extract 1µg of tyrosine per 1min. at 30C for 30 min.

The method of hydrolysis

- 1. pH about 1.5~5.0; temp. about 30~50C.
- 2. Hydrolyzing until 40~70% of the solubility of 0.75M trichloroacetic acid.
- 3. Use heat or pH adjust to deactivate the proteinase, and centrifuge it to remove undisolved whey protein.
- 4. Dry the peptide solution, and collect the powder.
- 5. Reconstitute the powder.
- 6. Purify the tripeptide by using membrane, ion exchange, gel filtration, and extract the tripeptide with a reverse column of HPLC.
 - a. Adjust pH3.0~5.0, and adsorb tripeptide with ion exchange chromatography (resin: Sp-Toyopearl 550C).
 - b. Desorb the component by using NaCl solution from 0M to 0.5M as liner gradient, and collect high ACE inhibitor which comes out when NaCl solution is 0.4M~0.5M.
 - c. Purify the solution with Bio-Rad Biogel P-2 column and distilled water.
 - d. Use HPLC(ODS-120T, Toso, Inc.) to absorb the solution, and desorb the peptide with mixture of 0.1% trifluoroacetic acid (A)and 0.1% trifluoroacetic acid that of 50% is acetonitrile (B).
 - e. Use liner gradient method that the amount of (B) increases gradually from 0% to 100%, and there is high concentration of ACE inhibiting activity when the concentration of acetonitrile is 20 to 22%.
 - f. Repeat above procedure as many as it needs, and dry it.
 - g. Use protein sequencer-(PSQ-1 system, Shimazu Manufacture) to confirm that tripeptide consist of Leu-Lys-Pro.

The method of tripeptide synthesis

- -Use peptide synthesis instrument (Biolynx 4170, Falmashia, Sweden).
- -Condense Fmocoproline on polyamide resin and remove Fmoc group from it.
- -Condense Fmocelysine on free amino group, and remove Fmoc group.
- -Condense Fmoceleucine on it, and remove Fmoc group.
- -Remove the resin from the peptide by using 95% trifluoroacetic acid at room temp, and remove trifluoroacetic acid by vacuum.
- -Mix with 0.1N acetic acid, and extract Leu-Lys-Pro with HPLC(ODS-120T).

Conclusion

This ACE inhibitor can be used small amount for humans and animals to decrease blood pressure and prevent hypertension. The amount of it is depend on age, weight, gender, symptom, and kind of animal and humans. It is possible to take it orally or not and to mix with liquid or solid carrier or without. It can be mixed with other medicines, and can be form of tablets, powder, capsules, injections. Natural ACE inhibitor, L-Leu-L-Lys-L-Pro, can be added to food and animal feed.

<u>Example</u>

- -Dissolve 5g of whey protein (ALACEN 132, Nihon Protein, Inc.) into 100ml of 0.03N HCl, and add distilled water up to 200ml.
- -Adjust pH 2.0 with 1N HCl, and react with 5000units of pepsin (Sigma) at 37C for 15 hours.
- -Adjust pH 4.4 with 5N NaOh, and react with 1000 units of Alpaltic Proteinase (ProteinaseM, Amano Pharmaceutical, Inc.) at 45C for 5 hours.
- -Adjust pH 6.0 with 5N NaOh, and heat at 90C for 20 min. to deactivate enzyme and precipitate non-dissolve materials.
- -Cool down the liquid temp., and centrifuge it at 10000G for 20 min. to remove solid materials.
- -Collect the supernatant, and freeze dry to get 4.0g of powder.
- -Dissolve 500mg of the powder into 50ml of 5mM acetic acid buffer, and adjust pH 3.5 with 1N HCl.
- -Run this solution into the column, 16mmX200mm, with 40ml of SP-Toyopearl 550C (Toso, Inc.) at 1.0ml per min. of flow, and run 120ml of NaCl solution by using liner gradient method from 0M to 0.5M at 1ml per min. of flow.
- -Collect high fraction of ACE inhibitor at 0.4~0.5M NaOh, and use the column, 16mmX1000mm, with 200ml of Biogel P-2 (Bio-Rad) at 0,33ml per min. of flow to filter.
- -Run distilled water to collect eluent.
- Use HPLC(ODS-120T, Toso, Inc.) to absorb the solution, and desorb the peptide with mixture of 0.1% trifluoroacetic acid (A)and 0.1% trifluoroacetic acid that of 50% is acetonitrile (B).
- -Use liner gradient method that the amount of (B) increases gradually from 0% to 100%, and there is high concentration of ACE inhibiting activity when the concentration of acetonitrile is 20 to 22%.
- -Repeat above procedure, and dry it.
- -Collect 1200µg of powder, and use protein sequencer (PSQ-1system, Shimazu Manufacture) to confirm that tripeptide consist of H•L-Leu-L-Lys-L-Pro•OH.

The method of ACE inhibiting activity for the peptide

- -Mix $50\mu l$ of the peptide solution with $20\mu l$ of ACE solution (mix 1 unit of ACE from rabbit lung of Sigma with $5\pi l$ of distilled water), and heat at 37C for 5 min.
- -Add the substrate (5mM Hip-His-Leu: pH8.3), and heat at 37C for 30 min., and add 1ml of 0.3M NaOh to stop the reaction.
- -Add 100µl of orthophthalate aldehyde, fluorescent tester, and react at room temp. for 10 min.
- -Add 200ml of 3N HCl, and dilute 50 times with distilled water.
- -After 30 min., measure the fluorescent intensity of the solution (A) at 300 μ m of excited wave and 490 μ m of fluorescent wave, and the fluorescent intensity of control sample without the peptide is (B).
- -Inhibiting activity formula: B-A/B
- -Adjust the activity to 50% by changing the concentration of the peptide.

(ACE inhibiting activity (IC50))

Peptide	IC50(μM)
H•L-Leu-L-Lys-L-Pro•OH	2.2
bradykinin potentiater B	6.4
bradykinin potentiater C	29.0

-The result shows that the small amount of the ACE inhibiting peptide invented can reach to IC50 that means high ACE inhibiting activity.

(Effect of invention)

- This ACE inhibitor can be used small amount to decrease blood pressure and prevent hypertension. It is easy to take it orally or not, because this white powder can be dissolve in water, etc.
- ACE inhibitor, Leu-Lys-Pro, is simple structure, 3-amino acid chain; therefore, it is also easy to produce this peptide by synthesis and to absorb into body.

Applicant: Nisshin Flour Product Patent attorney: Chika Takagi